

# Polysaccharide-synthesizing Glycosyltransferases and Carbohydrate Binding Modules: the case of Starch Synthase III

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**Abstract:** Glycosyltransferases (GTs) are a ubiquitous group of enzymes that catalyze the transfer of sugar moieties from activated donor molecules to specific acceptor molecules, forming glycosidic bonds. Nucleotide-sugars, lipid phosphate sugars and phosphate sugars can act as activated donor molecules while acceptor substrates involve carbohydrates, proteins, lipids, DNA and also, numerous small molecules (i. e. antibiotics, flavonols, steroids). GTs enzyme families are very ancient. They are founded in all the three domains of life and display three different folds (named GT-A, GT-B and GT-C) which are a variant of a common  $\alpha/\beta$  scaffold. In addition, several GTs contain an associated non-catalytic carbohydrate binding module (CBM) that could be critical for enzyme activity.

This work reviews the current knowledge on the GTs structures and functions and highlights those enzymes that contain CBMs, particularly starch binding domains (SBDs). In addition, we also focus on *A. thaliana* starch synthase III enzyme, from the GT-5 family. This protein has a GT-B fold, and contains in its N-terminal region three in tandem SBDs, which are essential for the regulation of enzyme activity.

**Keywords:** Carbohydrate binding modules, enzymes, glycosyltransferases, starch, starch binding domains, starch synthase.

## INTRODUCTION

### Structure and classification of glycosyltransferases

Glycosyltransferases (GTs) are defined as the enzymes that transfer an activated glycosyl donor, with a phosphate leaving group, to an acceptor molecule [1]. The most common sugar substrates are nucleotide sugars such as uridine nucleotides (i.e. UDPGlc, UDPGal, UDPGalNAc); adenine nucleotides (ADPGlc); and citidine and guanidine nucleotides such as CDPGlc and GDPMan, however, GTs could also use other donors such as nucleosides monophosphates and lipid phosphate sugars. Nucleotide sugars GTs are often referred to as “Leloir enzymes” in honor to L.F. Leloir, who discovered the sugar nucleotides and their role in the biosynthesis of carbohydrates [1, 2].

The acceptor substrates could be a wide range of biomolecules, including lipids, proteins and oligosaccharides, thus synthesizing glycolipids, glycoproteins and glycoconjugates. Based on the type of sugar that they transfer, GTs are grouped into several families (i.e. glucosyltransferases, galactosyltransferases, sialyltransferases, etc.).

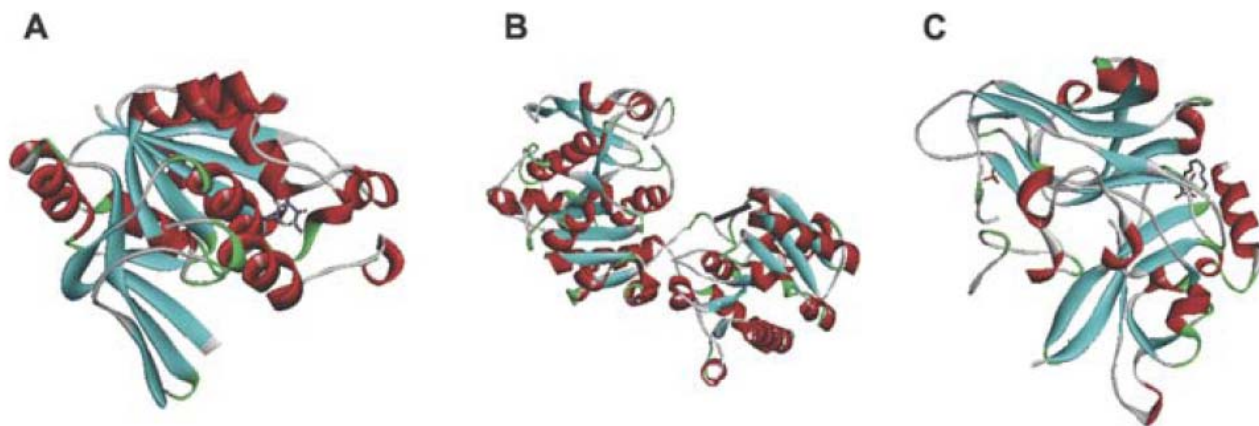
In addition, GTs can also be classified as either retaining or inverting enzymes according to the anomeric configuration of the substrates and reaction products. This stereochemically classification is the result of the different mechanisms used by the two groups or enzymes [3].

Retaining GTs operate via an internal return S<sub>N</sub>i-like mechanism with the retention of stereochemistry at the anomeric reaction center of the donor substrate; whereas inverting GTs most likely follow a single displacement mechanism where the acceptor substrate performs a nucleophilic attack at carbon C-1 of the sugar donor [1].

In the case of GTs, the enzymes within the same family catalyze the reaction with the same stereoselectivity. However, despite the fact that many GTs recognize identical donor or acceptor substrates, there is a very low sequence identity among the enzymes from different families [4].

It has been described that GTs adopt two different folds named GT-A and GT-B (Fig. 1) [5, 6]. The GT-A fold comprises two dissimilar domains, one for nucleotide binding and the other for the binding of the acceptor molecule, and was described for example for the SpsA protein, a nucleotide-diphosphosugar transferase from *Bacillus subtilis* involved in the synthesis of the spore coat and belonging to the GT-2 family (Fig. 1A) [7, 8]. The GT-B fold was described initially for the bacteriophage T4 beta-glucosyltransferase (EC 2.4.1.27) which catalyses the transfer of glucose from UDPGlc to cytosine bases in T4 duplex DNA [9]. Furthermore, the GT-B fold is present in other families such as the GT-3 and GT-5 families including about 90 glycogen or starch synthases from different organisms. The first protein whose structure was described in this group of GTs was the glycogen synthase from *Agrobacterium tumefaciens* (Fig. 1B) [10]. It was reported that the overall fold and the architecture of the active site of the protein are remarkably similar to those of glycogen phosphorylase [10].

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**Figure 1.** Ribbon structures showing the folds described for GTs: (A) GT-A fold represented by the SpsA protein, a nucleotide-diphosphosugar transferase from *Bacillus subtilis* (PDB ID 1H7L); (B) GT-B fold, represented by the glycogen synthase from *Agrobacterium tumefaciens* (PDB ID 1RZU); (C) GT-C fold, from EmbC, the arabinosyltransferase from *Mycobacterium tuberculosis* (PDB ID 3PTY).  $\alpha$  helices are colored in red, whereas  $\beta$  strands are represented in cyan.

to those of glycogen phosphorylase [10]. Nevertheless, the overall fold of the protein does not dictate the stereochemistry of the reaction that it catalyzes. Coutinho *et al.* [11] described several examples where both, inverting and retaining glycosyltransferases belongs to the GT-A and GT-B fold classes.

A third fold was also described, the GT-C fold which is a predicted protein topology for integral membrane glycosyltransferases (Fig. 1) [12]. However, this fold was not experimentally verified until the characterization of the 3D structure of a soluble C-terminal domain of an oligosaccharyltransferase STT3 from *Pyrococcus furiosus* [13]. Recently it has also been described other GT-C type GT essential for the synthesis of arabinogalactan and lipoarabinogalactan polymers in *Mycobacterium tuberculosis*, the EmbC arabinosyltransferase (PDB ID 3PTY, Fig. 1C) [14]. The co-crystallization of EmbC with a di-arabinoside analogue suggest that this protein contains two different carbohydrate binding sites, associated with subdomains I and II, respectively. Furthermore, the subdomain II functions as a carbohydrate-binding module (CBM) [14].

To date, GTs have been classified into 94 distinct families based in sequence data, available at the Carbohydrate Active Enzymes database server (CAZy, <http://www.cazy.org/>) [15]. The last family, 94, was recently defined after the biochemical characterization of the GumI protein, a non-processive  $\beta$ -mannosyltransferase that uses GDPMan as glycosyl donor substrate, essential for xanthan polysaccharide synthesis [16].

However, GT families 36, 46 and 86 were recently extinguished. The report on the evolutionary, structural and mechanistic relationship of the chitobiose phosphorylase with GHs of the clan GH-L provided insights to merge of GT family 36 to the glycosyl hydrolase family 94 [17]. The lack of consistent data for the biochemical characterization of GT-46 members led to the extinction of such family while members of GT-86 were merged to GT-83 (CAZY, <http://www.cazy.org/>).

The first 3D structures described for GTs were those from a glucanotransferase, the cyclodextrin GT from *Bacillus circulans* (EC 2.4.1.19, PDB code 1CGT), in 1991 [18, 19]; and the  $\beta$ -GT from bacteriophage T4 mentioned above (EC 2.4.1.27, PDB code 1BGT) [9]. The  $\beta$ -GT belongs to the GT-63 family, whereas the cyclodextrin GT was classified into the glycoside hydrolase 13 family (GH-13) in the CAZY server. While only few structures of GTs were described about 20 years ago, at the present it have been reported the 3D structure of 124 proteins belonging to 40 different GT families (Table 1). In addition, there are 1457 structure hits in the protein data bank (<http://www.rcsb.org/>) corresponding to GTs (EC 2.4.x.x). Most of the 3D structures (about 1437) were solved by X-ray diffraction, whereas few structures were determined by solution NMR.

### Carbohydrate Binding Modules

Most of the enzymes that degrade, modify, or create glycosidic bonds share a common modular organization. Such disposition usually includes a catalytic domain typical for each enzyme and one or more carbohydrate binding domain (CBM) connected by a loosely-structured chain. CBMs were initially classified as cellulose binding domains (CBDs), based on the initial discovery of several modules with the ability to bind this polysaccharide [20, 21]. Nonetheless, there are other types of modules different from these that can bind carbohydrates different from cellulose. These findings created the need for a new classification of these peptides using a more inclusive terminology. Thus, a CBM is nowadays defined as the contiguous amino acid sequence in an active enzyme involved in carbohydrate metabolism, with the ability to bind carbohydrates [22]. CBMs contain between 30 and 200 amino acids and may exist as simple, double or triple repeats in a protein (there are rare instances of independent putative CBMs). These domains can occupy N or C-terminal positions and occasionally can be found in the middle of the polypeptide chain.

CBMs have been found in glycoside hydrolases, glycosyltransferases, polysaccharide lyases, as well as carbohydr-

Table 1. Structures Reported for Polysaccharide-synthesizing Glycosyltransferases (Until September 2012\*)

Family (total structures)	Protein	PDB code	Reference
GT-1 (22)	GtfA <i>Amycolatopsis orientalis</i>	1PN3	[51]
GT-1 (22)	GtfA <i>Amycolatopsis orientalis</i>	1PN3	[51]
GT-2 (6)	SpsA <i>Bacillus subtilis</i>	1H7L	[8]
GT-3 (1)	glycogen synthase 2 (Gsy2, <i>S. cerevisiae</i> )	3NB0	[52]
GT-4 (15)	sucrose synthase (Sus1, <i>A. thaliana</i> )	3S27	[53]
GT-5 (3)	glycogen synthase 1 (GlgA, <i>A. tumefaciens</i> )	1RZU	[10]
GT-6 (3)	$\alpha$ -1,3 galatyltransferase (GGTA1, <i>Bos taurus</i> )	1FG5	[54]
GT-7 (3)	$\beta$ -1,4 galactosyltransferase-7 (b4Galt-7, <i>D. melanogaster</i> )	3LW6	[55]
GT-8 (4)	glycogenin (Gyg1, <i>Homo sapiens</i> )	3Q4S	[56]
GT-9 (4)	ADP heptose LPS heptosyltransferase II (WaaF, <i>E. coli</i> )	1PSW	<a href="http://www.rcsb.org">http://www.rcsb.org</a>
GT-10 (1)	$\alpha$ -1,3 fucosyltransferase (FucT, <i>H. Pylori</i> )	2NZW	<a href="http://www.rcsb.org">http://www.rcsb.org</a>
GT-13 (1)	<i>N</i> -acetylglucosaminyltransferase I (GnT1, rabbit)	1FO8	[57]
GT-14 (1)	2 beta1,6- <i>N</i> -acetylglucosaminyltransferase (C2GnT-L, <i>M. musculus</i> )	2GAK	[58]
GT-15 (1)	$\alpha$ -1,2-mannosyltransferase (Kre2, <i>S. cerevisiae</i> )	1S4N	[59]
GT-20 (2)	$\alpha$ -trehalose-phosphate synthase (OtsA, <i>E. coli</i> )	1GZ5	[60]
GT-23 (2)	$\alpha$ -1,6- <i>L</i> -fucosyltransferase (NodZ, <i>Bradyrhizobium</i> sp.)	2HHC	[61]
GT-27 (3)	UDP-GalNAc:polypeptide $\alpha$ - <i>N</i> -acetyl galactosaminyltransferase-2 ( <i>H. sapiens</i> )	2FFU	[28]
GT-28 (1)	MurG, <i>E. coli</i>	1F0K	[62]
GT-29 (1)	$\alpha$ -2,3-sialyltransferase (ST3Gal I, <i>Sus scrofa</i> )	2WML	[63]
GT-30 (1)	Membrane GT (WaaA, <i>A. aeolicus</i> )	2XCI	[64]
GT-31 (1)	<i>O</i> -fucosylpeptide $\beta$ -1,3-GlcNAc transferase ( <i>M. musculus</i> )	2J0A	[65]
GT-35 (6)	glycogen phosphorylase (PygM, rabbit muscle)	1A8I	[66]
GT-41 (3)	UDP-Glc: protein <i>N</i> - $\beta$ -glucosyltransferase (HMW1C, <i>Actinobacillus pleuropneumoniae</i> )	3Q3E	[67]
GT-42 (2)	$\alpha$ -2,3-sialyltransferase (Cst-I, <i>C. jejuni</i> )	2P2V	[68]
GT-43 (3)	beta1,3-glucuronyltransferase I (GlcAT-I, <i>H. sapiens</i> )	1FGG	[69]
GT-44 (4)	Toxin A ( <i>Clostridium difficile</i> )	3SRZ	[70]
GT-51 (6)	Penicillin binding protein ( <i>A. baumannii</i> )	3UDF	[71]
GT-52 (1)	$\beta$ -galactosamide $\alpha$ -2,3-sialyltransferase (NST, <i>N. meningitidis</i> )	2YK4	[72]
GT-53 (1)	$\alpha$ -1,5-arabinosyltransferase C (EmbC, <i>M. tuberculosis</i> )	3PTY	[14]
GT-55 (2)	$\alpha$ -mannosyltransferase (MPG, <i>Pyrococcus horikoshii</i> )	2ZU7	<a href="http://www.rcsb.org">http://www.rcsb.org</a>
GT-63 (1)	DNA $\beta$ -glucosyltransferase (BGT, Bacteriophage T4)	1BGT	[9]
GT-64 (1)	$\alpha$ - <i>N</i> -acetylhexosaminyltransferase (Ext12, <i>M. musculus</i> )	1OMX	[73]
GT-65 (1)	GDP-Fuc: protein <i>O</i> -fucosyltransferase 1 (PoFut1, <i>C. elegans</i> )	3ZY2	[74]
GT-66 (4)	oligosaccharyltransferase (AglB, <i>Archaeoglobus fulgidus</i> )	3VGP	[75]
GT-68 (1)	GDP-L-Fucose: protein <i>O</i> -fucosyltransferase 2 (PoFUT2, <i>H. sapiens</i> )	4AP5	[76]
GT-70 (1)	$\beta$ -1,2-glucuronosyltransferase (GumK, <i>Xanthomonas campestris</i> )	2HY7	[77]
GT-72 (1)	DNA $\alpha$ -glucosyltransferase (AGT, Bacteriophage T4)	1XV5	[78]

(Table 1) Continued...

Family (total structures)	Protein	PDB code	Reference
GT-78 (1)	Mannosylglycerate synthase (Mgs, <i>Rhodothermus marinus</i> )	2BO4	[79]
GT-80 (4)	$\alpha$ -2,3/2,6-sialyltransferase 1 (St1, <i>Pasteurella multocida</i> )	2IHJ	[80]
GT-81 (3)	glucosyl-3-phosphoglycerate synthase (GpgS, <i>M. tuberculosis</i> )	3E25	[81]
GT-88 (2)	elongation factor 1A $\alpha$ -glucosyltransferase (Lgt1, <i>Legionella pneumophila</i> )	2WZG	[82]

\*One representative member of each family is described

rate esterases. The CBMs can increase the rate of catalysis by approximating the catalytic domain of the protein to its substrate or act as a scaffolding domain that organizes the catalytic subunits into a cohesive multi-enzymatic complex [22]. Thus, they have three general roles with respect to the function of their cognate catalytic modules: (i) a proximity effect, (ii) a targeting function and (iii) a disruptive function [23]. Currently, more than 300 putative sequences in over 50 different species have been identified and the binding domains have been classified in 64 families, based on their amino acid sequence, substrate binding specificity and structure at the CAZY server (<http://www.cazy.org/>) [22-26].

It is important to note that several 3-D structures of members from at least 23 representative families have been obtained, in some cases as protein-ligand complexes. Data from these structures shed light on the underlying mechanism for each interaction between a CBM and its substrate. CBMs from different families are structurally similar and their ability to bind carbohydrates can be attributed, at least partially, to several aromatic residues which assemble into a hydrophobic surface [22].

It is interesting to note that glycosyltransferases, the main topic of this review, are represented in 4 of the 64 CBM families already mentioned and are listed in Table 2. There are several polypeptide N-acetylgalactosaminyltransferases in the CBM13 family from different organisms including *C. elegans*, *B. Taurus*, *D. melanogaster*, *T. gondii* and *H. sapiens*. These GTs belong to GT-2 family and catalyze the transfer of an N-acetyl-D-galactosamine residue to a serine or threonine residue to glycosylate the receptor protein [27]. The x-ray crystal structure of the human enzyme (ppGalNAcT-2 (hT2)) is available and shows the dynamic association between the catalytic and lectin domains where CBMs are located [28].

The CBM20 family is one of the protein families where the starch binding function of these modules has been demonstrated in several cases (<http://www.cazy.org/>). This family include different GTs from Archea, such as an  $\alpha$ -cyclomaltodextrin glycosyltransferase [29]; several  $\beta$ -cyclodextrin glycosyltransferases from bacteria, which cyclizes part of the  $\alpha$ -1,4 glucan chain [30, 31] and two glucanotransferases, essential for maltose metabolism in plants and probably, playing a role in freezing tolerance [32].

CBM48 family is represented by GT from bacteria and eucaryota. All examples involve branching enzymes that catalyzes the formation of  $\alpha$ -1,6-glucosidic bonds in bacterial glycogen and plant starch as well as in the synthesis of glycogen in mammals. One of the most characterized mem-

bers of this family is the starch branching enzyme I from *O. sativa* which its 3D structure was recently solved [33] (see also Table 2).

The last family is the CBM53 family, which currently has about 20 characterized entries in the CAZY database. All these entries correspond to starch synthase III (SSIII) belonging to higher plant species (*Arabidopsis thaliana*, *Zea mays*, *Solanum tuberosum*, *Phaseolus vulgaris*, *Vigna unguiculata* and *Oryza sativa*) and *Chlamydomonas reinhardtii*. It has been described that SSIII proteins contains three in tandem N-terminal Starch Binding Domains (SBDs) [34, 35].

The SBDs have acquired the evolutionary advantage to break the structure of their substrate when compared to the CBD, due to the presence of two binding sites for the polysaccharide [36]. So far, no CBD structure containing such binding sites has been reported. In view of this, such domains are unable to break down the polysaccharide structure as efficiently as the SBD does [36, 37]. Thus, SBDs are involved in the attachment of the catalytic domain to its polymeric substrate as well as in the disruption of the surface of the starch granule [38, 39].

Currently, SBDs are categorized into the ten following CBM families: (i) CBM20, i.e., the C-terminal SBD from *Aspergillus niger* glucoamylase; (ii) CBM21, located at the N-terminal domain in amylases; (iii) CBM25, containing one (i.e.,  $\beta$ -amylase from *B. circulans*) or two (i.e., *Bacillus* sp.  $\alpha$ -amylase) modules; (iv) CBM26, mostly organized in tandem repeats (i.e., C-terminal domains from *Lactobacillus manihottivorans*  $\alpha$ -amylase); (v) CBM34, present in the N-terminal domains of neopullulanase, maltogenic amylase, and cyclomaltodextrinase; (vi) CBM41, N-terminal SBDs, present mostly in bacterial pullulanases; (vii) CBM45, originating from eukaryotic proteins from the plant kingdom (i.e., N-terminal modules of  $\alpha$ -amylases and  $\alpha$ -glucan water dikinases); (viii) CBM48, modules with glycogen-binding function (including SBD from the GH13 pullulanase and regulatory domains of mammalian AMP-activated protein kinase); (ix) CBM53, SBD modules from SSIII; and, (x) CBM58, represented by *Bacteroides thetaiotaomicron* SusG protein that has been shown to bind maltoheptose [15, 24, 25, 40]. However, CBM20 and CBM21 families are proposed to form a common clan [23]. Analyses of three-dimensional structures from many of these families showed a characteristic SBD fold of a distorted  $\beta$ -barrel composed of several  $\beta$ -strands and with one or, more frequently, two distinct binding sites that exhibit site-dependent modes of carbohydrate binding [25, 26, 41-43].



**Table 2.** CBMs Families Present in Currently Characterized Polysaccharide-Synthesizing Glycosyltransferases

Family	Protein	PDB code	Reference
CBM13	UDP-GalNAc:polypeptide $\alpha$ -N- acetylgalactosaminyltransferase T2 (GalNAc-T2;ppGalNAcT2;GalNT2) ( <i>H. sapiens</i> )	2FFU	[28]
CBM20	$\beta$ -cyclodextrin glycosyltransferase ( <i>Bacillus circulans</i> )	--	[31]
CBM48	starch-branching enzyme 1 (Sbe1;Rbe1;BEI;Os06g0726400) ( <i>Oryza sativa Japonica Group</i> )	3AMK	[33]
CBM53	Starch synthase III (At1g11720) ( <i>A. thaliana</i> , <i>Chlamydomonas reinhardtii</i> 137C, <i>O. sativa Japonica Group</i> , <i>Pyricularia zeae</i> , <i>Solanum tuberosum</i> , <i>Vigna unguiculata</i> , <i>Zea mays</i> )	--	[35]

One example of each family is at least indicated. In case of various structures solved, the more recent is informed.

### Role of SBDs in Starch Synthase III

An outstanding example of the importance of the presence of CBMs, specifically SBDs, in the primary structures of a sugar-nucleotide GT enzyme is the SSIII from *Arabidopsis thaliana*. Through studies using null mutants from *A. thaliana* for the *AtSS3* locus, it has been postulated that SSIII could be a regulatory enzyme for the starch synthesis process, since these mutant lines displayed a higher starch accumulation in leaves during the day due to an apparent increase in biosynthetic rate [44]. Besides, SSIV has been proposed, along with SSIII, to be required at the onset of the starch granule synthesis [45, 46].

Members of the SS family display a high degree of sequence similarity in their C-terminal catalytic region. However, among the isoforms, the differences are mainly present at the N-terminal domain [47].

AtSSIII, with a total of 1025 amino acid residues, is a particular enzyme since it contains a putative N-terminal transit peptide followed by a 557-amino acid SSIII specific domain (SSIII-SD) and a C-terminal catalytic domain of 450 amino acids [34, 48]. We previously built a 3D structural model of the catalytic domain of AtSSIII using the structure of the glycogen synthase (GS, PDB ID 1RZU) from *A. tumefaciens* [48] (see also Fig. 1B). It is important to mention that the comparison between the model and GS crystal structure revealed that the global topology is almost the same, except for differences in some loops [10, 48]. Thus, the catalytic domain of AtSSIII, as well as the bacterial GSs share the same GT-B fold and were classified in the GT-5 family of glycosyltransferases.

The SSIII-SD contains three internal repeats that encode SBDs, named D1, D2 and D3. These SBDs were classified in the CBM-53 family, which currently has 89 members, mainly the SSIII from different plants and algae (including those from *A. thaliana*, *Oryza sativa*, *Ostreococcus lucimarinus*, *Ostreococcus tauri*, *Solanum tuberosum* and *Zea mays*, among others); and two glycoside hydrolases of *Thermococcus* sp. from the GT-57 family (<http://www.cazy.org/>).

Recently, we reported the role of the SDBs from AtSSIII by characterizing different forms of the enzyme lacking one, two or the three domains [35]. AtSSIII uses preferentially ADPGlc as glycosyl donor substrate, although UDPGlc can also be used [35]. When ADPGlc is the substrate, the presence of the SBDs confers distinctive properties to each en-

zyme isoform, increasing the apparent affinity and the  $V_{\max}$  for the oligosaccharide acceptor substrate (starch or glycogen). However, no substantial changes in the kinetic parameters for the polysaccharide were observed when UDPGlc was used as donor substrate. Under the acceptor polysaccharide saturating conditions, the presence of SBDs increases progressively the apparent affinity and  $V_{\max}$  for ADPGlc but not for UDPGlc [35]. Adsorption assays of the different isoforms to raw starch showed that SSIII-SD, containing three, two, or one SBD modules has an increased capacity to bind the polysaccharide depending on the number of SBD modules. The D23 protein, containing the second (D2) and third (D3) SBD modules is the one that makes the greatest contribution to polysaccharide binding [35]. Thus it is clearly demonstrated that SBD associate modules have a critical role, particularly the D23 domain, in the binding capacity to starch and in the regulation and catalysis of the GT activity of AtSSIII. Thus, the interaction of the N-terminal SSIII-SD, specifically the D2 and D3 domain (residues 316–344 and 495–535, respectively), with the catalytic domain showing GT activity, as well as the full integrity of the starch-binding capacity of the D2 domain, are involved in the modulation of AtSSIII [49].

Binding studies to raw starch and its components, amylose or amylopectin, show that the SBD binds preferentially to amylopectin, being the D1 domain primarily responsible for this selective binding. The D2 domain contains two polysaccharide binding sites which include amino acid residues Y394 (which is part of the binding site 1) and W366 (in the binding site 2) acting cooperatively with the D1 domain in the binding process. It was described that mutations in these residues also affect the kinetic parameters for the polysaccharide substrate of AtSSIII [50].

It is worth mentioning that most of SBDs are found in degrading starch and glycogen microbial enzymes. However, our work was the first report on the existence of an SBD in a synthesizing enzyme (AtSSIII) and, the first experimental evidence of its starch binding capacity which allowed to classify these modules within the CBM53 family [35, 49, 50].

### CONCLUSIONS

Thus, until present it is possible to define 91 (formerly 94 because 3 have been deleted or reclassified) GT families based on sequence homology. Despite of the very low se-

quence identity among the enzymes from different families, all enzymes remarkably converge in only three folds named GT-A, GT-B and GT-C. Hence, other properties must contribute to the huge functional plasticity of GTs; such as a large structural variability in the acceptor binding domain, the presence of CBMs, that could be involved in the modulation of the GT catalytic activity, as well as the possibility of supramolecular organization of GTs within the cell.

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## CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.

## ABBREVIATIONS

GT	=	(glycosyltransferase)
CBM	=	(carbohydrate binding module)
SBD	=	(starch binding domain)
SSIII	=	(starch synthase III)

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